(12)

EUROPEAN PATENT APPLICATION published in accordance with Art. 158(3) EPC

(21) Application number: 92910150.9

2 Date of filing: 15.05.92

(65) International application number: PCT/JP92/00633

(g) International publication number: WO 92/20739 (26.11.92 92/29) (i) Int. Cl.5: **C08L** 3/00, C08L 23/26, C08L 29/04, B32B 9/00, B32B 27/28

Priority: 15.05.91 JP 140948/91 01.07.91 JP 188016/91

1.07.91 JP 188016/91

27.08.91 JP 242535/91

30.09.91 JP 280749/91

30.09.91 JP 280750/91

- Date of publication of application: 05.05.93 Bulletin 93/18
- Designated Contracting States:
 DE FR GB IT

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◉ BIODEGRADABLE RESIN COMPOSITION AND LAMINATED STRUCTURE MADE THEREFROM.

 \odot A biodegradable resin composition comprising a saponified ethylene/vinyl acetate c⁻polymer and a starch, wherein the copolymer has an ethylene content of 20 to 60 mole %, a degree of saponification of the vinyl acetate units of 90 mole % or above, a melt flow index of 1 to 100 g/10 min at 210 °C under a 2,160 g load, and an η_{50} to η_{5} ratio of 0.5 to 4, wherein η_{50} is the m It viscosity of the sample kept at 250 °C for 60 min and η_{5} is the one at 250 °C for 5 min. The layer of this composition can form a laminated structure with another material such as a photodegradable polymer.

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TECHNICAL FIELD

The present invention relates to a biodegradable resin composition comprising a saponified ethylene – vinyl acetate copolymer and a starch – based macromolecular substance and to a laminate utilizing the same.

BACKGROUND ART

(Prior art)

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Molded plastic products such as plastic packaging films and containers, agricultural plastic films, binding tapes and so on are sometimes disposed of in ways other than legitimate routes of disposal. Because these products retain their shapes for years, they present serious environmental problems when discarded as they are. Therefore, it is highly desirable that any shaped products that are liable to be discarded in the open field, farmlands, rivers, etc. be those which will be rapidly decomposed by microorganisms in the soil or water.

Against this background, much development work has been undertaken to develop biodegradable compositions for the fabrication of such biodegradable shaped products. Referring to research and development along this line, the following patent applications have been filed in the technical field of bio – degradable or disintegratable compositions each comprising a saponified ethylene – vinyl acetate copolymer (i.e. ethylene – vinyl alcohol copolymer) and a starch – based macromolecular substance.

Japanese Patent Kokai Publication No. 3 - 31333

The official gazette of Japanese Patent Kokai Publication No. 3 – 31333 discloses a polymer composition for the manufacture of shaped biodegradable plastic products which comprises an ethylene – vinyl alcohol copolymer and a modified starch wherein the ethylene content of said ethylene – vinyl alcohol copolymer is 10 to 90 weight % and the melt flow index thereof is 2 to 50. The melt flow index here is the value determined at 230 °C under a load of 2.16 kg. The same gazette states that the preferred parameters of the ethylene – vinyl alcohol copolymer are as follows.

- o Intrinsic viscosity [η] (in dimethyl sulfoxide, 30°C): 0.5 0.9, preferably 0.60 0.80
- o Molecular weight distribution Mw/Mn (GPC in tetrahydrofuran): 1.3 4
- o Melting point: <180°C, preferably 160 170°C
- o Degree of hydrolysis: 90 99.9%

The same gazette states that, as the plasticizer, a polyol such as polyethylene glycol having a molecular weight of 200 to 4000, ethylene glycol, propylene glycol, sorbitol, or preferably glycerin can be employed.

Japanese Patent Kokai Publication No. 2-14228

The official gazette of Japanese Patent Kokai Publication No. 2-14228 discloses a polymer blend obtainable from a melt comprising a water-containing starch hydrolyzate and at least one substantially water-insoluble synthetic thermoplastic polymer. Claim 4 in the Scope of Demand for Patent of this gazette is directed to the use, as the thermoplastic polymer, of ethylene-vinyl acetate copolymer, ethylene-vinyl alcohol copolymer, ethylene-methyl acrylate copolymer, ethylene-methyl acrylate copolymer, ABS copolymer, styrene-acrylonitrile copolymer or polyacetal, thus referring to ethylene-vinyl alcohol copolymer, but there is no description about its ethylene content, nor does the specification include an example employing an ethylene-vinyl alcohol copolymer.

This gazette mentions, as the plasticizer, a variety of plasticizers such as low molecular weight polyalkylene oxides, e.g. polyethylene glycol, polypropylene glycol, polyethylenepropylene glycol, etc., low molecular weight organic plasticizers such as glycerol, glycerol monoacetate, diacetate or triacetate, etc., as well as propylene glycol, sorbitol, sodium diethylsulfosuccinate, triethyl citrate, tributyl citrate and so on.

Japanese Patent Kokai Publication No. 3-24101

The official gazette of Japanes Patent Kokai Publication No. 3-24101 discloses a method of producing a modified starch composition for the manufacture of shaped biodegradable plastic products which comprises blending starch with a high boiling plasticizer and a modifying agent selected from among

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urea, alkaline earth or alkali metal hydroxides and mixtures thereof at a temperature of 120 to 170 °C which is lower than the boiling point of the plasticizer for a sufficient time to modify the starch. Claim 5 in the Scope of Demand for Patent of this gazette states that the blending is performed in the presence of up to 15 weight % of the total weight of the mixture of an ethylene – acrylic acid copolymer and/or a polyvinyl alcohol. Furthermore, immediately preceding the Examples, the same gazette states that this modified starch composition is suited for blending with an ethylene – vinyl alcohol copolymer and an example is given as Example 5 in which a composition comprising 60 weight % of modified starch in the pellet form and 40 weight % of an ethylene – vinyl alcohol copolymer (Kuralene R20 (trademark)) is extruded. However, there is no specific reference to the ethylene content.

This gazette mentions, as the preferred plasticizer, such species as glycerin, polyethylene glycol, preferably polyethylene glycol having a molecular weight of 200 to 4000, ethylene glycol, propylene glycol, sorbitol and so on.

(Problems that the Invention is to Solve)

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Any biodegradable/disintegratable composition comprising an ethylene – vinyl alcohol copolymer and a starch – based macromolecular substance contains a large amount of the starch – based macromolecular substance which is liable to undergo pyrolysis in melt – molding and the ethylene – vinyl alcohol copolymer contained is also one of the resins comparatively difficult to melt – mold among the ordinary thermoplastic resins. Therefore, in this binary system, molding troubles such as deposits of burned resin around the die, gelation, coloration on aging, fish eye, etc. tend to take place in long – run production. In order to supply a biodegraddable/disintegratable product at low cost and on a steady basis in such a binary system, these problems associated with long – run moldability must be overcome.

However, in the gazette of Japanese Patent Kokai Publication No. 3-31333 referred to above, the ethylene - vinyl alcohol copolymer is selected only with reference to the commonplace indicators such as ethylene content, degree of saponification, melt flow index, intrinsic viscosity (an indicator of molecular weight), melting point and molecular weight distribution and, in regard to the melt characteristics of resin, attention is paid only to the parameters of melting point and melt flow index. Check experiments of the Examples described in Japanese Patent Kokai Publication No. 3-31333 indicate that although early moldability is acceptable on the whole, a variety of defects develop in moldings on long-run molding. This is probably because they did not have a definite idea about the melt characteristics necessary for long-run molding.

The official gazette of Japanese Patent Kokai Publication No. 2-14228 shows no specific example in which an ethylene-vinyl alcohol copolymer might have been actually used and, of course, the long-run moldability with this copolymer is not discussed.

Example 5 in the official gazette of Japanese Patent Kokai Publication No. 3-24101 indicates an example in which a composition comprising 60 weight % of pelletized modified starch and 40 weight % of an ethylene-vinyl alcohol copolymer was extruded using an extruder but there is no discussion on the melt characteristics of the copolymer, nor is the subject of long-run moldability explored, irrespective of the use of the copolymer.

After all, the above official gazettes of Japanese Patent Kokai Publication Nos. 3-31333, 2-14228 and 3-24101 show a mere background technology concerning the biodegradable/disintegratable composition comprising a saponified ethylene-vinyl acetate copolymer (ethylene-vinyl alcohol copolymer) and a starch-based macromolecular substance and the long-run moldability of such a binary system remains to be elucidated yet.

Moreover, as mentioned above, any starch – based macromolecular substance is thermolabile and, when it is blended with a resin and subjected to melt – forming, undergoes pyrolysis at temperatures above about 180°C. On the other hand, the saponified ethylene – vinyl acetate copolymer is one of the thermoplastic resins which are difficult to melt – form. For example, the melting point of a highly saponified ethylene – vinyl acetate copolymer is not less than 180°C when its ethylene content is about 30 mole % or less. Therefore, a binary system comprising these two materials tends to cause molding troubles or defects in shaped products. Therefore, in commercial production, it is essential to incorporate a variety of plasticizers mentioned above prior to molding. How ver, the application of those plasticizers which are commonly used in the conventional process s to the binary system comprising a saponified ethylene – vinyl acetate copolymer and a starch – based macromolecular substance results in the problem of poor com – patibility, inadequacy in the clarity and flexibility of the film or shet formed therefrom, and changes in physical properties and contamination owing to bleeding of the plasticizer. Thus, improvements that would satisfy all of these requirements have been much awaited.

It is an object of the pr sent invention to provide a resin composition capable of providing a biodegradable or biodisintegratable shaped product.

It is a further object of the invention to improve the melt-moldability, particularly long-run moldability, of a binary system comprising a saponified ethylene-vinyl acetate copolymer and a starch-based macromolecular substance.

A further object of the invention is to provide a resin composition which, when molded into film or sheet with the aid of a plasticizer, insures satisfactory clarity and flexibility and is less liable to undergo changes in physical properties or fouling due to bleeding of the plasticizer.

A still further object of the invention is to provide said composition conducive to a reduced amount of the plasticizer required.

It is a further object of the invention to provide a laminate comprising a layer composed of said composition and a backing layer (particularly a substrate film composed of a photodegradable polymer).

DISCLOSURE OF INVENTION

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The biodegradable resin composition of the present invention comprises a saponified ethylene – vinyl acetate copolymer having an ethylene content of 20 to 60 mole %, a vinyl acetate saponification degree of not less than 90 mole %, a melt flow index of 1 – 100 g/10 min. as measured under a load of 2160 g at 210 °C and a melt viscosity ratio η_{60}/η_5 , between the melt viscosity η_{60} after 60 minutes at 250 °C, of 0.5 – 4 and a starch – based macromolecular substance.

The laminate of the present invention is a laminar structure comprising a layer comprosed of said biodegradable resin composition and a layer composed of a different material (particularly at least one photodegradable polymer selected from the group consisting of ethylene – CO copolymer, ethylene – vinyl ketone copolymer, polybutadiene and isoprene rubber or a polyolefin type photodegradable polymer containing at least one of said polymers).

The present invention is now described in detail.

Saponified ethylene - vinyl acetate copolymer

As the saponified ethylene – vinyl acetate copolymer, a copolymer having an ethylene content of 20 – 60 mole %, preferably 25 – 50 mole %, and a vinyl acetate saponification degree of not less than 90 mole %, preferably not less than 95 mole % is employed. If the ethylene content is less than 20 mole %, the melt – moldability of the composition is sacrificed. If 60 mole % is exceeded, the mechanical strength and gas barrier property of the product are impaired so that the practical utility of the composition is compromised. If the vinyl acetate saponification degree is less than 90 mole %, the mechanical strength will not be sufficient. Moreover, outside of the above – mentioned range, not only the physical characteristics of the product but also the compatibility of the polymer with the starch – based macromolecular substance is adversely affected.

As the saponified ethylene – vinyl acetate copolymer, two or more species varying in ethylene content can be employed in combination. An example is the use, in combination, of a saponified ethylene – vinyl acetate copolymer having an ethylene content of 40 – 60 mole % and a saponified ethylene – vinyl acetate copolymer having an ethylene content of 20 – 50 mole %, the ethylene content of the former being greater than that of the latter by at least 5 mole %, in a weight ratio of 70:30 through 30:70.

Within the limits not jeopardizing the object of the invention, the saponified ethylene – vinyl acetate copolymer of the above composition may have been modified by copolymerization with other copolymerizable monomers [other α – olefins, ethylenically unsaturated carboxylic acid compounds (acids, anhydrides, salts, esters, amides, nitriles, etc.), vinyl ethers, vinyl esters other than vinyl acetate, ethylenically unsaturated sulfonic acid compounds (acids, salts, etc.), oxyalkylene – containing monomers and so on] or "post – modified" by oxyalkylene – etherization, cyanoethylation, acetalization, urethanation,

Regarding the melt-characteristics of the saponified thylene-vinyl acetate copolymer having the above composition, it is first of all necessary that the copolymer should have a melt flow index of 1 to 100 g/10 min, Preferably 1 to 60 g/min, and more preferably 2 to 30 g/10 min, as measured under a load of 2160 g at 210 °C. If the melt flow index is outside of the above range, the polymer will be unqualified as a melt-molding grade.

Another essential melt - characteristic for the saponified ethylene - vinyl acetate copolymer for us in accordance with the invention is that the melt viscosity ratio η_{60}/η_{5} , i.e. the ratio of the melt viscosity η_{60} after 60 minutes at 250 °C to the melt viscosity η_{5} after 5 minutes at 250 °C, should be 0.5 to 4 and

preferably 0.8 to 2.5. This is an essential element of the present invention. If η_{60}/η_{5} is less than 0.5, that is to say the tendency to viscosity d crease exceeds the allowable range, long-run molding results in the d position of burned resin around the die, infiltration of gels, formation of fish eyes, coloration on aging and other product defects as well as molding instability in the manufacture of a strand or film. If η_{60}/η_{5} exceeds 4, that is to say the tendency toward viscosity gain exceeds the allowable range, long-run molding results in the deposition of burned resin around the die, infiltration of gels, formation of fish eyes and other product defects. Thus, in either case, the long-run molding operation is made difficult.

The melt viscosity mentioned above is the value determined with a Koka type flow tester using a 1 mm ϕ x 10 mm nozzle and a load of 10 kg/cm². The values η_5 , η_{60} are found by measuring the viscosities at the residence times in the cyclinder of 5 and 60 minutes at a temperature of 250 °C.

Such saponified ethylene – vinyl acetate copolymer varies its properties according to the polymerization method, saponification method, degree of polymerization, degree of saponification, treatment following saponification, drying conditions, additives, etc. and the method of production cannot be defined in general terms. Typically, however, it can be obtained by saponifying an ethylene – vinyl acetate copolymer having the above – defined ethylene content with alkali and purifying the resulting saponified ethylene – vinyl acetate copolymer powder, grains or pellets by washing. In this purification by washing, the resin is thoroughly washed with an aqueous solution of acid, especially of a weak acid, or a dilute aqueous solution of a strong acid or an acidic salt thereof and, if necessary, further rinsed to remove the acid attached to the resin, followed by drying.

The weak acid which can be used as above generally includes acids having pKa (25°C) values of not less than 3.5, such as acetic acid, propionic acid, glycolic acid, lactic acid, adipic acid, azelaic acid, glutaric acid, succinic acid, benzoic acid, isophthalic acid, terephthalic acid and so on. It is possible and preferable to further treat the resin with a dilute strong acid, e.g. an aqueous solution of an organic acid having a pKa (25°C) value of not more than 2.5, such as oxalic acid, maleic acid, etc., an inorganic acid such as phosphoric acid, sulfuric acid, nitric acid, hydrochloric acid, etc. or an acidic salt thereof before or after aqeuous rinse following said treatment with a weak acid.

When the η_{60}/η_{5} values of commercial saponified ethylene – vinyl acetate copolymers are investigated from the above standpoint of melt characteristics, they are found to vary a great deal from one showing a tendency toward viscosity gain to one showing a tendency toward viscosity drop, indicating that no attention has heretofore been paid to η_{60}/η_{5} .

The saponified ethylene – vinyl acetate copolymer can also be obtained as one having a water content of 20 to 60 weight % according to conditions of production. The compatibility of such a copolymer with the starch – based macromolecular substance is superior and can be smoothly melt – molded even without use of a plasticizer or with a reduced amount of the plasticizer. However, an excessive water content should be avoided, for otherwise foaming may take place in melt – molding to interfere with smooth molding or the surface characteristics of moldings will be adversely affected. Typically scuh a saponified ethylene – vinyl acetate copolymer can be prepared by adding water to a methanolic solution of the saponified ethylene – vinyl acetate copolymer obtained by alkali saponification of an ethylene – vinyl acetate copolymer to give a solution in the mixed solvent of water – methanol, introducing this solution into water for coagulation, pelletizing or crushing the coagulate, and purifying the same by washing. In the purification by washing, it is preferable, just as described before, to wash the resin thoroughly with acid, particularly an aqueous solution of a weak acid or a dilute aqueous solution of an acidic salt of a strong acid, removing the acid attached to the resin by rinsing, and drying the resin under conditions conducive to the desired water content.

Starch - based macromolecular substance

The starch – based macromolecular substance includes raw starches such as corn starch, potato starch, sweet potato starch, wheat starch, cassava starch, sago starch, tapioca starch, sorghum starch, rice starch, legume starch, arrowroot starch, bracken starch, Indian lotus starch, water chestnut starch, etc.; physically modified starch (α – starch, fractionated amylose, wet heat – treated starch, etc.), enzymatically modified starch (hydrolyzed dextrin, enzymatically degraded dextrin, amylose, etc.); chemically degraded and modifi d starch (acid – treated starch, hypochlorite – oxidized starch, dialdehyde starch, etc.); chemically modified starch derivatives (esterifi d starch, therized starch, cationized starch, crosslinked starch, etc.) and so on. Among said chemically modified starch derivatives, esterified starch includes acetic acid – esterified starch, succinic acid – esterified starch, nitric acid – sterified starch, phosphoric acid – esterified starch, vanthic acid – esterified starch, acetoacetic acid – esterified starch, etc. The etherized starch includes allyl – therized starch, methyl – etherized starch, etc. The cationized therized starch, hydroxyethyl – etherized starch, hydroxypropyl – etherized starch, etc. The cationized

starch includes the reaction product between starch and 2-diethylaminoethyl chloride and the reaction product between starch and 2,3-epoxypropyltrimetylammonium chloride, etc. The crosslinked starch includes formaldehyde-crosslinked starch, epichlorohydrin-crosslinked starch, phosphoric acid-crosslinked starch, acrolein-crosslinked starch and so on.

In the practice of the present invention the starch – based macromolecular substance can be used in hydrated state and the proper water content in that case may for example be 8 to 30 weight %. Then, coloration, burning, charring and contamination with foreign matter may be alleviated.

Proportions

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The ratio of saponified ethylene – vinyl acetate copolymer to starch – based macromolecular substance is preferably 80:20 through 20:80, w/w, and more preferably 70:30 through 30:70, w/w. If the proportion of starch – based macromolecular substance is too small, the biodegradability and disintegratability will be poor, while an excessive proportion of the substance detracts from the mechanical properties of moldings. However, depending on applications, some derivation from the above ratio can be tolerated.

Additives

While the resin composition of the present invention is essentially composed of said saponified cthylene – vinyl acetate copolymer and starch – based macromolecular substance, it may contain as necessary a variety of additives such as plasticizers (e.g. polyhydric alcohols, urea, etc.), resin components (e.g. ethylene copolymers and other polyolefins, hydrogenated styrene – butadiene rubber, polyurethane, polyamide, polyhydroxybutyrate, etc.), natural polymers other than those related to starch (polysaccharide type polymers, cellulosic polymers, proteinaceous polymers, etc.), heat stabilizers, diluents, autoxidants, fillers, lubricants, colorants, flame retardants, waterproofing agents, ultraviolet absorbers, crosslinking agents, fungicides, herbicides, antioxidants and so on. As starch modifiers, urea, alkaline earth or alkali metal hydroxides or mixtures thereof can also be added.

It is particularly instrumental to incorporate a polyhydric alcohol, such as glycerin, trimethyolpropane, pentaerythritol, ethylene glycol, diethylene glycol, triethylene glycol, propylene glycol, dipropylene glycol, etc. as well as urea. The amount of such plasticizer can be optionally selected.

In the practice of the invention, the polyhydric alcohol preferably has 2 to 4 hydroxyl groups and melts at a temperature not below 40°C, preferably between 40 and 150°C, and has a molecular weight of not more than 600, preferably between 100 and 300. Among examples of such polyhydric alcohol are dihydric alcohols such as 1,6 – hexanediol, 1,8 – octanediol, 1,9 – nonanediol, 1,10 – decanediol, neopentyl glycol, hydrobenzoin, 1,2 – cyclohexanedimethanol, etc.; trihydric alcohols such as trimethylolpropane, trimethylolethane, etc.; and tetrahydric alcohols such as pentaerythritol. Among them, 1,6 – hexanediol, neopentyl glycol, 1,4 – cyclohexanedimethanol, and trimethylolpropane are preferred from practical points of view. The proportion of such polyhydric alcohol is 5 to 50 parts by weight, preferably 10 to 40 parts by weight, based on 100 parts by weight of the sum of said saponified ethylene – vinyl acetate copolymer and starch – based macromolecular substance. If the proportion is less than 5 parts by weight, the effect will not be sufficient, while a proportion of more than 50 parts by weight will not insure a commensurate effect but rather detract from mechanical strength to make the composition unpractical.

Melt - molding technology

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As the melt - molding technology, extrusion molding (inclusive of blow molding) can be mainly utilized. However, calendering and injection molding processes can also be employed. The preferred extrusion molding process comprises feeding the saponified ethylene - vinyl acetate copolymer and the starch - based macromolecular substance either independently or concurrently to an extruder for melt - mixing and pelletization and feeding the resulting pellets to an extruder for molding to the desired shape. The melt - forming temperature in the latter stage is often set to a temperature not exceeding 180°C. When the starch - based macromolecular substance or the saponified ethylene - vinyl acetate copolymer is hydrous or when the molding operation is performed with addition of water, it is pref rable to reduce the water content of the extrudate to 5 w ight % or less using an extruder equipped with a vent means before feeding to the latter stage extruder for melt - molding.

When the melt-molding is carried out by extrusion molding, it is generally pref rable to set the resin temperature at the joint between the extruder and the die to 170°C or below and the shear rate of the melt passing through the die lip to not less than 100 sec⁻¹.

In molding, it is also possible to employ a special mixing mode such that the saponified ethylene – vinyl acetate copolymer and the starch – based macromolecular substance are independently melted beforehand and, then, blended in molten stat. According to this mode, there can b manufactured a shaped product of improved quality with reduced incidences of discoloration associated with thermal degradation, fish eyes, grains, gelation and burn. In this case, a plasticizer may be incorporated but its proportion need not be greater than, for example, 10 parts by weight relative to 100 parts by weight of the sum of saponified ethylene – vinyl acetate copolymer and starch – based macromolecular substance. Regarding the above mode in which the saponified ethylene – vinyl acetate copolymer melted by heating beforehand is blended with the starch – based macromolecular substance similarly melted beforehand, an exemplary process comprises melting the starch – based macromolecular substance in a twin – screw extruder in the first place and feeding the saponified ethylene – vinyl acetate copolymer from a side feed port of the same twin – screw extruder for mixing to give a homogeneous blend. For the manufacture of shaped products, it is preferable to pelletize the blend thus obtained and feed the pellets to an extruder for molding to the desired shape.

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Laminate

In accordance with the present invention, a layer composed of such a composition comprising the saponified ethylene - vinyl acetate copolymer and starch - based macromolecular substance can be lami - nated with a layer composed of a different substrate material. The different material mentioned just above may for example be a thermoplastic resin, kraft or other paper, woven cloth, nonwoven cloth, metal foil or wood board.

The thermoplastic resin includes, among others, polyolefin resin, vinyl alcohol resin, acrylic resin, polyamide resin, polyester resin, polyurethane resin, polystyrene resin, polyacetal resin, polycarbonate resin, diene type resin, cellulosic resin and so on. When, among these thermoplastic resins, particularly one or more photodegradable polymers selected from the group consisting of ethylene—CO copolymer, ethylene—vinyl ketone copolymer, polybutadine and isoprene rubber or a polyolefinic photodegradable polymer containing at least one of these resins as blended, or even a photodegradable polymer rendered photodegradable by incorporation of a transition metal or a photosensitizer is employed as said different material for lamination with the layer comprised of the composition of the invention which is per se biodegradable, the resulting laminate as a whole is naturally disintegratable. The laminate obtainable by using a paper, a natural cellulose—based woven or nonwoven cloth, a wooden board or the like is also biodegradable and disintegratable. Therefore, an increased industrial usefulness can then be attained.

The ethylene – CO copolymer can be obtained by copolymerizing ethylene with CO at high temperature and super – high pressure and the copolymerizing rate of CO is generally about 0.3 to 10 weight %. The ethylene – vinyl ketone copolymer includes ethylene – methyl vinyl ketone copolymer and ethylene – phenyl vinyl ketone copolymer, to name but a few. The copolymerizing proportion of vinyl ketone is generally about 0.1 to 10 mole % but may be higher. The polybutadiene includes thermoplastic 1,2 – polybutadiene, high cis – butadiene and low cis – butadiene rubbers, to name but a few. Particularly preferred is a thermoplastic 1,2 – polybutadiene with 90% of butadiene units forming 1,2 – bonds. The isoprene rubber includes natural rubber, isoprene rubber, trans – polyisoprene, etc. and, among them, natural rubber is particularly important.

It is also possible to employ a photodegradable polymer prepared by blending 3 to 20 parts by weight of any of said ethylene – CO copolymer, ethylene – methyl vinyl ketone copolymer, ethylene – phenyl vinyl ketone copolymer, 1,2 – polybutadiene, etc. with 100 parts by weight of a polyolefin such as a low – density polyethylene, linear low – density polyethylene, high – density polyethylene, polypropylene, ethylene – vinyl acetate polymer or the like. It is also possible to employ a photodegradable polymer whose photodeg – radability has been imparted by a transition metal salt or a photosensitizer.

As the transition metal salt, these can be employed salts of stearine, dialkylthiocarbamates, salicylal – dehyde, acetylacetone benzophenone oxime, etc. with iron(II), zinc(II), manganese(II), nickel(II), cerium(III), cobalt(II) or the like. The photosensitizer may for example be benzophenone, acetophenone or anth – raquinone.

For laminating a layer comprised of th above composition with a layer comprised of said different material, any of the known m thods such as dry lamination, extrusion, co-injection, multi-layer inflation, multi-layer T-di extrusion, multi-layer blow molding, multi-layer pipe extrusion, etc. can be invariably employed. As to the bonding mode of molten resin, whichever of "in-die" bonding and "out-of-die" bonding can be employed.

In manufacturing such a laminate by melt-molding, an adhesive can be employed with advantage. For the adhesive layer, a variety of adhesives such as polyolefin, polyester, acrylic and other adhesives can be

successfully utilized. Preferably, acid – modified polyolefins, and more preferably carboxylic acid (e.g. maleic anhydride) – modified polypropylene, carboxylic acid – modified ethylene – vinyl acetate copolymer adhesives are employed. Such an adhesive layer is generally used in a thickness of 5 to 50 µm. As the anchor coating agent for dry lamination, isocyanate, polyethyleneimine, polybutadiene, organotitanium and other coating agents can be advantageously employed and the coating thickness is generally 0.5 to 2 µm.

Other

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The photodegradable polymer mentioned hereinbefore may be incorporated in a composition compris – ing said saponified ethylene – vinyl acetate copolymer and starch – based macromolecular substance. The proportion of the photodegradable polymer for this purpose is generally 3 to 70 parts by weight based on 100 parts by weight of the sum of saponified ethylene – vinyl acetate copolymer and starch – based macromolecular substance.

5 Operation and Result

The present invention employs a saponified ethylene – vinyl acetate copolymer having a definite melt characteristic, that is to say one having the specified melt viscosity ratio η_{60}/η_{5} , where η_{60} means the melt viscosity after 60 minutes of standing at 250 °C and η_{5} means the melt viscosity after 5 minutes of standing at 250 °C, so that despite it being a binary system comprising a saponified ethylene – vinyl acetate copolymer and a starch – based macromolecular substance, the acceptable melt moldability and particularly long – run moldability which are the long – standing problems to be solved are remarkably improved. Therefore, the invention permits a stable supply of biodegradable shaped products in this binary system.

Furthermore, when a polyhydric alcohol containing 2 to 4 hydroxyl groups within the molecule and having a melting point of not lower than 40°C and a molecular weight of not more than 600 is employed as the plasticizer, molding compatibility is very satisfactory and films of excellent clarity and flexibility can be manufactured. Moreover, bleeding of the plasticizer can also be avoided.

When a hydrous saponified ethylene - vinyl acetate copolymer with a water content of 20 to 60 weight % is employed as said saponified ethylene - vinyl acetate copolymer, satisfactory moldability is assured even if addition of the plasticizer is omitted or the amount of the plasticizer is drastically reduced.

Furthermore, when the saponified ethylene - vinyl acetate copolymer and the starch - based macro - molecular substance are independently melted beforehand and blended in molten state, satisfactory melt - molding results can be obtained even with a reduced amount of the plasticizer.

Laminating a layer composed of the above composition with a substrate layer (particularly a layer composed of at least one photodegradable polymer selected from the group consisting of ethylene – CO copolymer, ethylene vinyl ketone copolymer, polybutadiene and isoprene rubber or a polyolefin – based photodegradable polymer blend containing at least one of said polymers) is useful in that it broadens the scope of applicability.

40 BEST MODE FOR CARRYING OUT THE INVENTION

The following examples are intended to describe the invention in further detail.

Examples 1 ~ 5, Comparative Examples 1 ~ 5

(Production of saponified ethylene - vinyl acetate copolymers)

To a methanolic solution of an ethylene – vinyl acetate copolymer having an ethylene content of 44 mole %, 38 mole %, 32 mole % or 29 mole % was added sodium hydroxide, as the catalyst, for primary saponification and the resulting preliminary hydrolysate was dispersed in water. Then, a further amount of sodium hydroxide was added for secondary saponification to obtain a saponified ethylene – vinyl acetate copolymer having a saponification degree of 99 mole %. This hydrolysate was rinsed with water and, where necessary, washed with a dilute aqueous solution of acetic acid or sodium phosphate, followed by drying. In this manner, saponified thylene – vinyl acetate copolymers having the characteristic values indicated hereinafter in Table 1 wer obtained.

(Melt - molding and evaluation of long - run moldability)

Each of the saponified ethyl n -vinyl ac tate copolymers obtained above was fed to a twin-screw extruder via its main feeder at a rate of 3 kg/hr. At the same time, a mixture of 100 parts by weight of starch (corn starch) and 30 parts by weight of glycerin was fed to the above twin-screw extruder via its side feeder at a rate of 4 kg/hr. The strand extruded through the die was pelletized with a pelletizer to give pelllets 3 mm in diameter and 3 mm in length. The temperature settings of the twin-screw extruder were as follows.

C ₁	C ₂	C₃	C₄	C₅	Н	D
200°C	170°C	160°C	160°C	160°C	160°C	160°C

This pelletizing operation was continued for 3 consecutive days (1 or 2 days in Comparative Exam – ples) and the incidences of deposition of burned resin near the discharge port of the twin – screw extruder, infiltration of gels into the pellet, and the degree of pellet discoloration were investigated. After completion of the above continuous molding, the twin – screw extruder was disassembled and inspected for residues of degradation products.

Then, the above pellets (those obtained up to 3 hours after the beginning of pelletization) were fed to a single – screw extruder for repelletizing and this cycle was repeated for a total of 6 times. The final pellets were fed to a single – screw extruder equipped with a T – die to give a 30 μ m thick film and the infiltration of gels into the film and the incidence of fish eye were investigated. Moreover, the extruder used for the production of film was disassembled and inspected for internal residues of degradation products.

In the above procedure, the single – screw extruder for pelletization and the single – screw extruder for film formation were set to the following temperatures.

Single - s	screw extru	der for pell	etization			
C ₁	C ₂	C₃	C ₄	C₅	н	D
130°C	160°C	160°C	160°C	160°C	160°C	160°C

Single -	screw extru	der for film	formation						
C ₁	C ₁ C ₂ C ₃ C ₄ C ₅ H D								
130°C	160°C	160°C	160°C	160°C	160°C	160°C			

Results

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The results of evaluation of long-run molding performance are shown below in Table 1. The characteristic values of the saponified ethylene-vinyl acetate copolymers used are also presented in Table 1

40 45	35		30		25	20	15	10	5	
					Table	. 1				
			Example	эте			Com	Comparative Example	ample	
	1	2	3	4	s	1	2	3	4	5
Characteristic valu	values of EVOH									
Et content (mole %)		44	38	32	53	44	38	38	32	59
SV (mole %) MFI (g/10min) · η _{ξΩ} /η ₅	99 12 1.2	99 12 0.9	99 25 2.2	99 3 2.5	99 8 2.0	99 12 4.9	99 8 5,5	99 25 0.1	99 3 5.2	99 8 0.3
moldab	ility with a twin-screw extruder	:win-scre	w extr	ıder						
Number of days of continuous molding	м _	е	m	٣	m	- .	7	7		2
Burnt resin deposits	ts None	None	None	Slight	None	Abundant	Abundant	Abundant	Abundant	Abundant
Gels	None	None	None	None	None	Abundant	Abundant	Obvious	Abundant	Obvious
Coloration on aging	g None	Minimal None	None	None	None	Minimal	Minimal	Obvious	None	Obv1ous
Degradation product residues	t None	None	None	Minimal	None	Abundant	Abundant	Abundant	Abundant	Obvious
(Disassembling inspection)	spection)									
Overall evaluation	0	٥	0	0	o	×	×	×	×	٧
Film formation after	9	pelletizing cycles	cles							
Gels	None	None	None	None	None	Abundant	Abundant	Abundant	Abundant	Obvious
Fish eyes	None	None	None	None	None	Abundant	Abundant	Abundant	Abundant	Abundant
Degradation product residues	t None	None	None	Minimal	None	Abundant	Abundant	Abundant	Abundant	Abundant
(Disassembling inspection) Overall evaluation	spection)	0	0	0	o	×	×	×	×	×

(Note) The symbols used in Table 1 have the following meanings.

EVOH: saponified ethylene-vinyl acetate copolymer

Et content: ethylene content

SV: saponification degree

MFI: melt flow index

 η_{60}/η_5 : melt viscosity ratio

(Note) In Table 1, the degrees of burnt resin deposit, gel infiltration, coloration on aging and fish eye were respectively evaluated on the 5-grade scale of none, minimal, slight, obvious and abundant.

The overall evaluation was made on the 3-grade scale of o (good), Δ (fair) and x (poor).

30 Examples 6 ~ 10

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(Production of saponified ethylene - vinyl acetate copolymers)

Each of ethylene - vinyl acetate copolymers varying in ethylene content was saponified, rinsed, washed with a dilute solution of acetic acid and a dilute aqueous solution of phosphoric acid, further rinsed and dried in the conventional manner.

The compositions, melt flow indices (MFI) and melt viscosity ratios of the resulting saponified ethylene – vinyl acetate copolymers are shown below in Table 2.

40 (Compound pelletization)

A Henschel mixer was charged with pellets of a saponified ethylene – vinyl acetate copolymer obtained above, raw starch (corn starch with a water content of 12 weight %, an example of the starch – based macromolecular substance), and a plasticizer in the proportions and combination shown in Table 2. After mixing, the mixture was fed to a twin – screw extruder equipped with vent means and melt – extruded into a strand and, at the same time, pelletized with a pelletizer to give a pelleted resin composition. The temperature settings of the twin – screw extruder were as shown below.

Screw diameter 30 mm

L/D 30

50 Screw RPM 150 rpm (unidirectional rotation)

Vent Open vent

Die Two nozzles, 5 mm in diameter

Tempera	ture setting	s				
C ₁	C ₂	C ₃	C4	C₅	Н	D
100°C	140°C	160°C	160°C	160°C	160°C	160°C

The resulting pellets were hot - pressed under the conditions of 150 °C x 50 kgf/cm² x 5 minutes to mold a 1 mm - thick plate. To evaluate the clarity of the plate, the total light transmittance of the hot - pressed plate was measured in accordance with JIS K7105. (Film production)

The above compound pellets were fed to a single – screw extruder equipped with a T – die to mold a 50 μ m – thick film. The film – forming conditions of the single – screw extruder were as follows.

Screw diameter

40 mm

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Screw RPM

50 rpm (unidirectional rotation)

Screw configuration

Full flight

Compression ratio

3.0

T - die

Coat hanger type

Die width

450 mm

Die lip opening

0.5 mm

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Tempera	ture conditi	ons			
C ₁	C ₂	C3	C ₄	Н	D
140°C	160°C	160°C	160°C	160°C	160°C

(Bleeding test)

The above film was sandwiched between strips of filter paper and allowed to stand at 20 °C and 65% RH. The staining of the filter paper due to bleeding of the plasticizer was investigated by washing the filter paper after standing with methyl alcohol and measuring the infrared absorption spectrum of a concentrate of the washing.

35 (Biodegradability)

The above film was cut to B5 size (25.8 mm x 18.2 mm) and buried in a compost – soil mixture. After 6 months, the film was taken out and visually inspected and evaluated according to the following criteria.

- a: Only residual fragments can be confirmed and no shape of the original film is retained.
- b: The original shape is retained but a multiplicity of cracks and openings are observed all over.

Results

The results as well as the formulations are shown in Table 2.

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Table 2

		Exar	nple		
	6	7	8	9	10
Characteristic values	of EVOH				
Et content (mole %)	44.2	49.0	38.2	28.9	44.1
SV (mole %)	99.5	99.2	99.4	99.3	96.0
MFI (g/10 min)	12	15	25	8	13
η ₆₀ /η ₅	0.9	2.6	2.2	2.0	0.9
Plasticizer	NPG	CHD	HD	TMP	PE
Molecular weight	104	144	118	134	136
Melting point (°C)	128-130	45-50	43	59	260
Formula (parts)					
EVOH	40	35	45	50	55
Raw starch	60	65	55	50	45
Plasticizer	20	15	25	30	40
Pelletizability	Good	Good	Good	Good	Goo
Total light					
transmittance (%)	75	78	82	76	74
Modulus in tension					
(kgf/cm²)	6300	4800	5100	6600	900
Film formability	Good	Good	Good	Good	Goo
Bleeding	None	None	None	None	Non
Biodegradability	a	a	a	a	a

(Note) The abbreviations in the plasticizer column mean the following.

NPG: neopentyl glycol

CHD: 1,4-cyclohexanedimethanol

HD: 1,6-hexanediol

TMP: trimethylolpropane

PE: pentaerythritol

Reference Examples 1 ~ 2, Comparative Examples 6 ~ 7

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The procedures of Examples 6 ~ 10 were repeated except that the compounds indicated below in Table 3 were used as the plasticizer. The results are set forth in Table 3.

Table 3

e= 😁 .

		Compara Example	tive
1	2	6	7
s of EVOH 44.2 99.5 12 0.9	49.0 99.2 15 2.6	38.2. 99.4 25 0.1	28.9 99.3 8 0.3
Gly 92 18	L-PEG 300 -15~6	H-PEG 4000 53~56	SB 182 97
40 60 20	35 65 15	45 55 25	50 50 30
Good	Good	Good	Good
80	78	0	80
5600 Good Observed b	5200 Good Observed b	12000 Poor -	30000 Poor - -
	Examp 1 5 of EVOH 44.2 99.5 12 0.9 Gly 92 18 40 60 20 Good 80 5600 Good Observed	S of EVOH 44.2 49.0 99.5 99.2 12 15 0.9 2.6 Gly L-PEG 92 300 18 -15~6 40 35 60 65 20 15 Good Good 80 78 5600 5200 Good Good Observed Observed	Example Example 1 2 6 s of EVOH 44.2 49.0 38.2 99.5 99.2 99.4 12 15 25 0.9 2.6 0.1 Gly L-PEG H-PEG 92 300 4000 18 -15~6 53~56 40 35 45 60 65 55 20 15 25 Good Good Good 80 78 0 5600 5200 12000 Good Good Poor Observed Observed —

(Note) The abbreviations in the plasticizer column mean the following.

Gly: glycerin

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L-PEG: low molecular weight polyethylene glycol

H-PEG: high molecular weight polyethylene glycol

SB: sorbitol

(Note) In the pelletizability column, "poor" means a difficulty in take-up of the strand.

In the film formability column, "poor" means a difficulty in the formation of film.

Therefore, the bleeding and biodegradability parameters in Comparative Examples 6 and 7 were not determined.

Examples 11 ~ 15

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(Production of saponified ethylene - vinyl acetate copolymers)

To a methanolic solution of an ethylene – vinyl acetate copolymers varying in ethylene content was added a methanolic solution of sodium hydroxide as the catalyst and a continuous saponification reaction was conducted in a reactor. The resulting solution of the saponified copolymer in methanol was diluted with water to prepare a mixed solution in methanol and water and the dilution was discharged from a nozzle into water at a temperature not exceeding 20 °C. Thereupon the solution became coagulated in a strand form with a diameter of about 3 mm. The coagulate was taken out from the water and cut.

The coagulated strand was pelletized with a pelletizer, rinsed to remove sodium acetate from the polymer, washed with a dilute aqueous solution of acetic acid and further with a dilute aqueous solution of phosphoric acid and dried under varying drying conditions to adjust to the target water content.

The characteristic values, water contents, and melting points (DSC peak temperatures) in hydrous state of the resulting hydrous saponified ethylene – vinyl acetate copolymers are shown in Table 4.

22 (Pelletization of compound)

A Henschel mixer was charged with pellets of the above hydrous ethylene – vinyl acetate copolymer and raw starch (corn starch with a water content of 12 weight %) in the proportions and combination indicated in Table 4. After mixing, the mixture was fed to a twin – screw extruder with vent means and melt – extruded in the form of a strand and pelletized with a pelletizer. The operating parameters of the twin – screw extruder were as follows.

Screw diameter 30 mm 1/D 30 Screw RPM 150 rpm Vent Open vent

Die Two nozzles, 3.0 mm in diameter

Temper	ature settin	ıgs				
C ₁	C ₂	C ₃	C ₄	C₅	Н	D
50°C	110°C	120°C	120°C	120°C	120°C	120°C

40 (Melt-molding)

The compound pellets obtained as above were fed to an injection molding machine with a cramping pressure of 100 tons and injection – molded under the following conditions to fabricate a sheet (flat) sized 100 mm x 1.00 mm x 1.5 mm (thickness), a No. 1 tensile testpiece according to JIS K – 7113, and a flexure testpiece according to JIS K – 2703.

Injection pressure 200 kg/cm²

Injection speed High pressure setting

Mold A flat sheet mold of fan gate type, side gate No. 1 Tensile testpiece mold and

flexure testpiece mold

50 Mold temperature 40°C

Tempera	ture conditions		
C ₁	C ₂	C₃	Nozzle
100°C	110-140°C	110-140°C	110-140°C

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(Results)

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The results as well as the conditions used are shown in Table 4.

Table 4

		Exa	nple		
	11	12	13	14	15
Characteristic values	of EVOH	-			20.0
Et content (mole %)	44.0	44.0	44.0	38.0	29.0
SV (mole %)	99.5	99.5	99.5	99.5	99.5
MFI (g/10 min)	12	12	12	25	8
η_{60}/η_{5}	0.9	0.9	0.9	2.2	2.0
Water content (wt%)	53	53	26	30	55
Melting point in					
hydrous state (°C)	96	96	100	98	93
Pormula (parts)	150	100	100	100	160
Hydrous EVOH	100	100	100	100	100
Raw starch	100				
Pelletizability	Good	Good	Good	Good	Goo
Pellet composition (wt%)				2.4
EVOH	33	26	40	38	34
Raw starch	42	50	47	48	41
Water	25	24	13	14	25
Injection molda-					0
bility of pellet	Good	Good	Good	Good	Goo
Mechanical strengths		· · · · · · · · · · · · · · · · · · ·			
Tentile strength			350	360	260
(kg/cm²)	240	300	40	35	40
Elongation (%)	50	40	40	د د	40
Flexural strength	220	280	350	350	250
(kg/cm ²)					
Bleeding	None	None	None	None	None
Biodegradability (di	sintegrata	ability)		_	•
Six months	b	þ	Ъ	р	р
One year	a	a	a	а	а

(Note) In the pelletizability column, "good" means satisfactory blending and good takeup of the strand.

The mechanical strengths were measured at 20 $^{\circ}$ C in accordance with JIS K-7113 (tensile test) and 50 JIS K-2703 (flexure test).

For the evaluation of biodegradability (disintegratability), the flat sheet samples obtained as above were buried in a compost-soil mixture and 6 months and one year later were taken out and visually examined and scored on the following 3-grade scale.

- a: Not I ss than 50% by volum of the sample has lost shape.
- 55 b: The nitre surface of the sample has been corrugated and the four dges are also indented, indicating that disintegration is in progress.
 - c: Recessed spots are locally found on the surface and edges of the sample

E ...

Film production Examples 1 ~ 4

(Production of hydrous saponified ethylene - vinyl acetate copolymers)

The hydrous saponified ethylene - vinyl acetate copolymers indicated in Table 5 were prepared in the same manner as Examples 11~ 15.

(Production of films from compositions)

A Henschel mixer was charged with pellets of the above hydrous ethylene – vinyl acetate copolymer, raw starch (corn starch with a water content of 12 weight %) and, as necessary, glycerin in the proportions and combinations shown in Table 5. The mixture was then fed to a twin – screw extruder equipped with vent means and melt – extruded into a strand at 120°C and pelletized with a pelletizer.

The resulting pellets were fed to a single-screw extruder and extruded through a T-die at a cylinder temperature of not higher than 140°C and a die temperature of not higher than 140°C to prepare a 35 μ m-thick film. The film was passed through a hot-current circulating drying furnace at 150°C to adjust the water content of the film to not more than 7%.

(Production of films)

The materials mentioned in each Example were fed in the indicated proportions to a single-screw extruder and extruded from a T-die at a cylinder temperature of 200 to 230°C and a die temperature of 200 to 230°C to prepare a 50 μ m-thick film.

(Conditions and Results)

Pelletizability ratings, compositions and film-forming properties are shown in Table 5.

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Table 5

	Film	producti	on Examp	ole
	1	2	3	4
Characteristic values	of EVOH			
Et content (mole %)	44.2	27.8	38.2	46.8
SV (mole %)	99.5	99.5	99.4	99.2
MFI (g/10 min)	12	8	25	14
	0.9	2.0	2.2	1.0
η ₆₀ /η ₅ Water content (wt%) Melting point in	26	40	35	40
hydrous state (°C)	100	95	97	93
Formula (parts)				
Hydrous EVOH	67.6	108.3	84.6	
Raw starch	56.8	39.8	51.1	73.
Glycerin	-	-	<u>-</u>	19
Pelletizability	Good	Good	Good	Good
Pellet composition (w	t %)			
EVOH	42.1	46.6	42.2	25.6
Raw starch	42.1	25.1	34.6	47.4
Water	15.8	28.3	23.2	20.2
Glycerin	-	-	_	6.8
Dry weight ratio of				
raw starch to EVOH	50/50	35/65	55/45	65/35
Film-forming				
property	Good	Good	Good	Good
Film code number	F-1	F-2	F-3	F-4

(Fabrication of laminates)

Examples 16 ~ 19

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Using an isocyanate anchor coating agent (D), the above film (F-1), (F-2), (F-3) or (F-4) was dry-laminated with the following substrate film (C-1), (C-2), (C-3) or (C-4) to fabricate a laminate with the layer thicknesses (μ m) of (F)/(D)/(C) = 35/1/50.

- (C-1): an ethylene CO copolymer film
- (C-2): a low-density polyethylene film blended with 10 weight % of an ethylene-methyl vinyl ketone (5 mole %) copolymer
- (C-3): a high-density polyethylene film blended with 10 weight % of a thermoplastic 1,2-polybutadiene
- (C-4): a linear low-density polyethylene film containing 0.5 weight % of ferric stearate. The laminar structures and measured data are shown in Table 6.

Table 6

	Example			
	16	17	18	19
Laminar structure				
Film (F)			F-3	
Base film (C)	C-1	C-2	C-3	C-4
Oxygen permeability				
$(cc/m^2 \cdot 24 \text{ hr} \cdot \text{atm})$				
25°C, 30% RH	0.5	0.3	0.4	1.5
25°C, 75% RH	1.5	1.0	1.7	7.3
Interlayer bond strength				•
(g/25 mm wide)	850	830	760	530
Natural disintegratability				
on soil	Yes	Yes	Yes	Yes
Oli BOTT				

(Note) Oxygen permeability was measured in accordance with ASTM D 3985 at 25°C, 30% RH and 75% RH using Modern Control's OX-TRAN 100.

The interlayer bond strength was measured between film layer (F) and base film layer (C) in accordance with JIS Z = 0237.

As to disintegratability, a film sample 1 m x 1 m was set with its film layer (F) in contact with the ground surface and the film was visually inspected after 6 months and one year. As a result, the films according to Examples 16 ~ 19 showed formation of many cracks after 6 months and had disintegrated completely out of shape by the end of one year.

35 Example 20

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The above film (F-1) was laminated with a 30 μ m-thick kraft paper by hot-pressing and the physical properties of the laminate were determined. The oxygen permeability was 0.5 cc/m²•24 hr•atm at 25°C, 30% RH and 1.6 cc/m²•24 hr•atm at 25°C, 75% RH. The interlayer bond strength could not be measured because of breakage of the kraft paper during determination. As to disintegratability, many cracks and holes were found after 6 months and no shape was retained after 1 year.

Comparative Example 8

On a 50 µm - thick polyethylene film surfaced with an anchor coating material was formed a 35 µm - thick starch layer by the casting method using raw starch (corn starch with a water content of 12 weight %). The oxygen permeability of this laminate film was 2.0 cc/m²•24 hr•atm at 25°C, 30% RH and 70 cc/m²•24 hr•atm at 25°C, 75% RH. As to interlayer bond strength, peeling occurred and the starch film was easily broken at 0 g/25 mm width.

Comparative Example 9

A Henschel mixer was charged with 80 parts by weight of a low – density polyethylene with a melt index of 6 g/10 min. (Mitsubishi Kas i Corporation, F – 161) and 20 parts by weight of raw starch (corn starch with a water content of 12 weight %) and after blending, the mixture was fed to an extruder for pelletization. The pellets were fed to an extruder and melt – extruded through a T – die at a cylinder temperature of 190°C and a die temperature of 180°C to provide a 30 μ m – thick film. When the proportion of raw starch was increased over the above value, no film could be produced.

Using an isocyanate anchor coating agent (D), the film obtained above was dry – laminated with a 30 μ m – thick polyethyl ne film and the physical properties of the laminate were determined.

The oxygen permeability of this laminate film was not less than 1000 cm/m²•24 hr•atm at 25°C, 30% RH and not less than 1000 cm/m²•24 hr•atm at 25°C, 75% RH. The interlayer bond strength was 850 g/25 mm width.

Example 21

(Production of saponified ethylene - vinyl acetate copolymers)

An ethylene – vinyl acetate copolymer was saponified and washed in the same manner as Examples 11 – 15 and dried to give the saponified ethylene – vinyl acetate copolymers indicated below in Table 7.

Table 7

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	E-1	E-2	E-3
Characteristic values of EVOH			
Et content (mole %)	29.2	38.8	43.3
SV (mole %)	99.5	99.4	99.5
Melting point (* C)	186	178	167
MFI (g/10 min)	8.2	25.4	12.1
750/75	1.9	2.2	0.9

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(Compound pelletization)

Starch and glycerin were admixed and the mixture was fed in a predetermined amount through a measuring feeder to a twin-screw extruder, in which it was melted by heating at 170°C. Then, a saponified ethylene-vinyl acetate copolymer prepared as shown in Table 7 was fed in a predetermined amount through a measuring feeder to a single-screw extruder and melted by heating at 210°C. The melt was fed to the above twin-screw extruder via its side feeding port. The two charges were blended until a homogeneous blend was obtained. The formulas are shown hereinafter in Table 8. The extruder parameters were as follows.

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Twin - screw extruder					
Screw diameter		30 mm			
L/D		30			
Screw RPM		120 rpm			
Extrusion temperatures	C ₁ C ₂ C ₃ C ₄ C ₅ Adapter Die	70°C 160°C 170°C 170°C 170°C 170°C			
(Side feeder: C ₃)					

Single - screw extruder				
Screw diameter 30 mm				
L/D	24			
Screw RPM		100 rpm		
Extrusion temperatures	C ₁ C ₂ C ₃ Joint	210°C 200°C 210°C 210°C		

(Production of films)

The compound pellets obtained by 12-hr continuous extrusion as externally supplemented with an aggregate (0.3 part of ethylenebisstearoidamide) was fed to a single-screw extruder equipped with a T-die to prepare a 50 μ m-thick film. The film-forming conditions at the single-screw extruder were as follows.

Screw inside diameter	40 mm
l r∖d	28
T-die	Coat hanger type
Die width	450 mm
Die lip opening	0.5 mm
Extrusion temperature	140 - 180°C

Table 8

	Example						
	21	22	23	24	25	26	27
Formula		-					
Twin-screw extruder Starch	70	120	120	200	120	200	300
(water content 12.8%) Glycerin	10	20	10	10	10	10	20
Single-screw extruder EVOH	E-1 100	E-1 100	E-2 100	E-2 100	E-3 100	E-3 100	E-3 100
Evaluation		_					
Processability Surface of strand Thermal coloration of pellet	Smooth Light yellow						
Quality Degree of granular structure	Good						
Tensile strength (kg/cm²) Elongation (%)	420 300	350 280	380 270	370 200	370 250	350 210	250 150

(Note) Processability was evaluated in terms of the surface condition of the strand obtained after 15 hours of continuous extrusion and the thermal coloration of the pellets. The surface condition of the strand follows a course toward roughening and the thermal coloration of pellets progresses in the order of light yellow (normal) to yellow to brown.

Quality was assessed in terms of the degree of entry of grains (degraded resin gels, burnt resin grains, charred resin grains, and unfused gels) sized 1 mm or larger per 30 cm x 30 cm of the film prepared from the pellets obtained after 15 hours of operation (≤5 good, 6−20, some, >20 abundant) and the tensile strength and elongation (as measured at a pulling speed of 500 mm/min, a chuck−to−chuck distance of 50 mm and a reference line − to − line distance of 20 mm) of the film.

INDUSTRIAL APPLICABILITY

Shaped products obtainable from the biodegradable composition of the invention find application in a variety of fields such as packaging materials (film, sheet, bottle, cup, tray, can carrier, agricultural materials (agricultural film, bundling tape, etc.) and civil materials (diaper backing sheet, shopping bag, garbage bag, etc.).

5 Claims

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 A biodegradable resin composition comprising a saponified ethylene - vinyl acetate copolymer having an ethylene content of 20 to 60 mole %, a vinyl acetate saponification degree of not less than 90 mole

%, a melt flow index of 1 to 100 g/10 min as measured under a load of 2160 g at 210 °C, and a melt viscosity ratio η_{60}/η_{5} of 0.5 to 4 where η_{60} means the melt viscosity after 60 minutes of standing at 250 °C and the η_{5} means the melt viscosity after 5 minutes of standing at 250 °C and a starch – based macromolecular substance.

- The biodegradable resin composition of claim 1 wherein the saponified ethylene vinyl acetate copolymer and the starch – based macromolecular substance occur in a weight ratio of 80:20 through 20:80.
- 70 3. The biodegradable resin composition of claim 1 wherein the saponified ethylene vinyl acetate copolymer is a hydrous saponified ethylene vinyl acetate copolymer having a water content of 20 to 60 weight %.
- 4. The biodegradable resin composition of claim 1 which contains a plasticizer in addition to said saponified ethylene vinyl acetate copolymer and starch based macromolecular substance.
 - 5. The biodegradable resin composition of claim 1 wherein the plasticizer is a polyhydric alcohol containing 2 to 4 hydroxyl groups and having a melting point of not lower than 40°C and a molecular weight of not more than 600.
 - 6. The biodegradable resin composition of claim 1 as produced by melting said saponified ethylene vinyl acetate copolymer and said starch based macromolecular substance independently and blending them in molten state.
- 25 7. A laminate comprising a layer composed of a biodegradable resin composition comprising a saponified ethylene vinyl acetate copolymer having an ethylene content of 20 to 60 mole %, a vinyl acetate saponification degree of not less than 90 mole %, a melt flow index of 1 to 100 g/10 min as measured under a load of 2160 g at 210° C, and a melt viscosity ratio η₆₀/η₅ of 0.5 to 4 where η₅₀ means the melt viscosity after 60 minutes of standing at 250° C and η₅ means the melt viscosity after 5 minutes of standing at 250° C and a starch based macromolecular substance and a substrate material of a different kind.
 - 8. The laminate of claim 7 wherein said different substrate material is a layer of at least one photodeg radable polymer selected from the group consisting of ethylene CO copolymer, ethylene vinyl ketone copolymer, polybutadiene and isoprene rubber or a polyolefinic photodegradable polymer containing at least one of said polymers as blended.

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INTERNATIONAL SEARCH REPORT

international Application No PCT/JP92/00633

		international Application No PCI	/3732/00033
	IFICATION OF SUBJECT MATTER (If several classification (IPC) or to both Nation		
int.	. C1 ⁵ C08L3/00, C08L23/26, B32B9/00, B32B27/28	CU8L29/U4,	
II FIELDS	BJZB9/UU, BJZBZ1/Z8		
	Minimum Document	ation Searched?	
lessification	on System (C	lassification Symbols	
IPO	C08L3/00, C08L23/26,	C08L29/04,	
	B32B9/00, B32B27/28		
	Documentation Searched other the to the Extent that such Documents a		
III. DOCU	MENTS CONSIDERED TO BE RELEVANT	·	
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į	Responsabilita Limitarta) February 12, 1991 (12. 02		
{	Claim (Family: none)	,,	
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	Claim, lines 19 to 20, up	per right column,	
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Y	JP, A, 56-14544 (The Nippe	on Synthetic	1-6
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* Special	categories of cited documents: 14	"T" later document published after t	he international filling date or
	ument defining the general state of the art which is not sidered to be of particular relevance	priority date and not in conflict w understand the principle or theor	ith the application but cited to ry underlying the invention
"E" earl	ier document but published on or after the international	"X" document of particular relevance be considered novel or cannot	
"L" doc	g date rument which may throw doubts on priority claim(s) or	"Y" document of particular relevance	
which cita	ch is cited to establish the publication date of another tion or other special reason (as specified)	be considered to involve an inverse is combined with one or more	ntive step when the documen
	tument referring to an oral disclosure, use, exhibition or an oral means	combination being obvious to a t	person skilled in the art
"P" doc	sument published prior to the international filing date but	"E" document member of the same (patent family
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	e Actual Completion of the International Search	Date of Mailing of this International S	Search Report
Jul	y 9, 1992 (09. 07. 92)	August 4, 1992 (
	nal Searching Authority	Signature of Authorized Officer	
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